derived from tissues which do not normally express P-glycoprotein but in which P-glycoprotein expression increases during the development of the cancer. One example is chronic myelogenous leukemia, which when it goes into blast crisis, expresses more P-glycoprotein irrespective of the previous treatment history (Gottesman, M.M. Cancer Research, 53:747-754 (1993)).

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On page 5, replace the paragraph starting on line 8 with the following:

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In another aspect, the invention includes a liposome composition for administration of a therapeutic compound to the cytoplasm of a cell characterized by increased expression of the MDR1 gene. The liposomes are composed of vesicle-forming lipids and include a vesicle forming lipid derivatized with a hydrophilic polymer chain having a free distal end. A folate ligand is attached to the free distal end of at least a portion of the hydrophilic polymer chains, and a therapeutic agent entrapped in the liposomes.

<u>In the Claims</u>: Please replace claim 1 with the following:

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 (Amended) A method of administering a therapeutic compound to a multi-drug resistant cell expressing Pglycoprotein, comprising

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preparing a conjugate composed of (i) a carrier; (ii) a folate ligand attached to the carrier; and (iii) a therapeutic agent associated with the carrier; and

administering the conjugate to a subject;

whereby said administering is effective to achieve accumulation of said therapeutic agent in said cell.